(FILE 'HOME' ENTERED AT 11:33:02 ON 28 NOV 2001)

FILE 'CAPLUS' ENTERED AT 11:33:16 ON 28 NOV 2001

	FILE 'CAPL'	JS, MEDLINE, EMBASE, BIOSIS' ENTERED AT 11:39:31 ON 28 NOV 2001
L1	36	S TNF ANTAGONIST (P) (ETANERCEPT OR INFLIXIMIB OR ANTI-TNF ANTI
L2	1	S L1 AND (SPINAL INJURY OR ALZHEIMER? DISEASE OR PALSY OR SPINA
L3	0	S L1 AND (LOCALIZED TREATMENT OR INTRALEISONAL? OR PERILEISON?)
L4	5	S L1 AND (SUBCUTANEOUS OR INTRAMUCULAR OR TRANSEPIDERMAL OR PAR
L5	5	DUP REM L4 (0 DUPLICATES REMOVED)
L6	120	S TNF (P) (SPINAL INJURY OR ALZHEIMER? DISEASE OR PALSY OR SPIN
L7	28	S L6 AND TNF (5A) (INHIBIT? OR REDUC? OR SUPPRESS? OR DOWNREGU
L8	16	DUP REM L7 (12 DUPLICATES REMOVED)
L9	1	S L8 AND (ETANERCEPT OR INFLIXIMIB OR MONOCLONAL ANTIBODY)

Refs: 56

ISSN: 0146-0404 CODEN: IOVSDA

COUNTRY: DOCUMENT TYPE:

United States Journal; Article

Ophthalmology FILE SEGMENT: 012

Clinical Biochemistry 029

English LANGUAGE: English SUMMARY LANGUAGE:

PURPOSE. The cytokine TNF.alpha. is a strong modulator of AB trabecular meshwork (TM) matrix metalloproteinase (MMP) and tissue inhibitor (TIMP) expression. Studies were conducted to identify signal-transduction pathways involved. METHODS. Porcine TM cells were treated with TNF.alpha., and MMP and TIMP levels were evaluated by zymography and Western immunoblot. Inhibitors and activators of several signal-transduction pathways were. . . were evaluated. PKC isoform down-regulation and additional inhibition profiles were used to refine the involvement pattern of different isoforms. RESULTS. TNF.alpha. treatment increased MMP-1, -3, and -9 and TIMP-1 expression, whereas MMP-2 expression was not affected and TIMP-2 expression decreased. Agents. . . modulate protein kinase A (PKA) or inhibit phosphatidylinositol 3-kinase (PI3K) had minimal effects on trabecular MMP or TIMP induction by TNF.alpha., whereas several agents that modulate PKC activity were effective. Trabecular cells expressed several PKC isoforms, which exhibited distinctive subcellular localization. TNF.alpha. treatment triggered some PKC isoform translocations. Exposure of trabecular cells to TNF.alpha. for 72 hours differentially downregulated several PKC isoforms. Treatment with a phorbol mitogen that stimulates most PKC isoforms produced strong increases in these MMPs. TNF.alpha.'s effects on MMP and TIMP expression were completely blocked by only one PKC inhibitor. CONCLUSIONS. The PKA and P13K pathways appear not to be involved directly in transducing this TNF.alpha. signal, but at least one isoform of PKC seems to be required. Based on the inhibitor profiles and the downregulation. signal. Unraveling the remaining steps in this and in additional related TM signal-transduction pathways may provide targets for developing improved glaucoma treatments.

DUPLICATE 1 MEDLINE L8ANSWER 5 OF 16

ACCESSION NUMBER:

DOCUMENT NUMBER:

2001423004

MEDLINE

PubMed ID: 11303144

Aging and proinflammatory cytokines. TITLE: Bruunsgaard H; Pedersen M; Pedersen B K AUTHOR:

Department of Infectious Diseases, H:S, Rigshospitalet, CORPORATE SOURCE:

University of Copenhagen, Denmark.

CURRENT OPINION IN HEMATOLOGY, (2001 May) 8 (3) 131-6. SOURCE:

Ref: 49

Journal code: CNO; 9430802. ISSN: 1065-6251.

United States PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

English LANGUAGE:

Priority Journals FILE SEGMENT:

200107 ENTRY MONTH:

Entered STN: 20010730 ENTRY DATE:

> Last Updated on STN: 20010730 Entered Medline: 20010726

Aging is associated with increased inflammatory activity reflected by AB increased circulating levels of TNF-alpha, IL-6, cytokine antagonists and acute phase proteins in vivo: Epidemiologic studies suggest that chronic low-grade inflammation in aging promotes an atherogenic profile and is related to age-associated disorders (eg, Alzheimer disease, atherosclerosis, type 2 diabetes, etc.) and enhanced mortality risk. Accordingly, a dysregulated production of inflammatory cytokines has an important role. . .

IN-PROCESS 2001548991 ACCESSION NUMBER:

21479537 PubMed ID: 11596043 DOCUMENT NUMBER:

Nerve injury proximal or distal to the DRG induces similar TITLE:

> spinal glial activation and selective cytokine expression but differential behavioral responses to pharmacologic

treatment.

Winkelstein B A; Rutkowski M D; Sweitzer S M; Pahl J L; AUTHOR:

DeLeo J A

Department of Anesthesiology, Dartmouth-Hitchcock Medical CORPORATE SOURCE:

Center, Lebanon, NH 03756.

JOURNAL OF COMPARATIVE NEUROLOGY, (2001 Oct 15) 439 (2) SOURCE:

127-39.

Journal code: HUV; 0406041. ISSN: 0021-9967.

United States PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

IN-PROCESS; NONINDEXED; Priority Journals

ENTRY DATE:

FILE SEGMENT:

Entered STN: 20011015

Last Updated on STN: 20011015

. . (3) responding to pharmacologic interventions. Rats received AB either an L5 spinal nerve transection distal to the DRG or an L5 nerve root injury proximal to the DRG.

Comparative studies assessed behavioral nociceptive responses, spinal cytokine mRNA and protein expression, and glial activation after injury. In separate studies, intrathecal pharmacologic interventions by using selective cytokine antagonists (interleukin-1 [IL-1] receptor

antagonist and soluble tumor necrosis factor [TNF]

receptor) and a global immunosuppressant (leflunomide) were performed to determine their relative effectiveness in these injury paradigms. Behavioral responses assessed. . . of persistent pain, suggesting that behavioral testing may not be a sensitive measure of injury. Spinal IL-1beta, IL-6, IL-10, and TNF mRNA and IL-6 protein were significantly elevated in both injuries. The overall magnitude of expression and temporal patterns were similar. . . for both injuries. In contrast, the pharmacologic treatments were more effective in alleviating mechanical allodynia for peripheral nerve injury than

nerve root injury, suggesting that nerve root injury elicits a more robust,

centrally mediated response than peripheral nerve injury. Overall, these data implicate alternate nociceptive mechanisms in these. . .

ANSWER 7 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS L8

2001:193568 BIOSIS ACCESSION NUMBER: PREV200100193568 DOCUMENT NUMBER:

Cytokine production consequent to T cell-microglia TITLE:

interaction: The PMA/IFNgamma-treated U937 cells display

similarities to human microglia.

Chabot, Sophie; Charlet, Danielle; Wilson, Tammy L.; Yong, AUTHOR(S):

V. Wee (1)

(1) Departments of Oncology and Clinical Neurosciences, CORPORATE SOURCE:

University of Calgary, 3330 Hospital Drive, NW, Calgary,

AB, T2N 4N1: vyong@ucalgary.ca Canada

Journal of Neuroscience Methods, (15 February, 2001) Vol. SOURCE:

105, No. 2, pp. 111-120. print.

ISSN: 0165-0270.

Article DOCUMENT TYPE: English LANGUAGE: English SUMMARY LANGUAGE:

where activated T cells, regardless of specificities, come into contact with microglia; these disorders include multiple sclerosis, trauma, stroke and Alzheimers disease. A model cell line would facilitate studies of the engagement between T cells and human adult microglia, since the latter. . . line shows similarities to microglia in its interaction with activated T lymphocytes, in that the production of tumor necrosis factor (TNF)-alpha, interleukin (IL)-4, IL-10 and IL-12 is induced. Morphological features and mechanisms of cytokine production resemble those observed in microglia-T cell

TNF-alpha and IL-10 levels, while anti-CD23 inhibits IL-10 only in U937-T cell interactions. We propose that PMA/IFNgamma-treated U937 cells can serve. . .

L8 ANSWER 8 OF 16 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:56156 CAPLUS

DOCUMENT NUMBER:

135:209814

TITLE:

Downregulation of Microglial Activation by Apolipoprotein E and ApoE-Mimetic Peptides

AUTHOR(S):

Laskowitz, D. T.; Thekdi, A. D.; Thekdi, S. D.; Han, S. K. D.; Myers, J. K.; Pizzo, S. V.; Bennett, E. R. Department of Medicine (Neurology), Duke University

CORPORATE SOURCE:

Medical Center, Durham, NC, 27710, USA Exp. Neurol. (2001), 167(1), 74-85

SOURCE:

CODEN: EXNEAC; ISSN: 0014-4886
Academic Press

DOCUMENT TYPE:

PUBLISHER:

Journal English

LANGUAGE:
REFERENCE COUNT:

72

REFERENCE(S):

(2) Avila, E; J Biol Chem 1982, V257, P5900 CAPLUS

(3) Barger, S; Nature 1997, V388, P878 CAPLUS

(4) Bellosta, S; J Biol Chem 1995, V270, P27063 CAPLUS

(6) Chen, Y; Neuroscience 1997, V80, P1255 CAPLUS (7) Clay, M; Biochemistry 1995, V34, P11142 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

AB Apolipoprotein E plays an important role in recovery from acute brain injury and risk of developing Alzheimer's disease. We demonstrate that biol. relevant concns. of apoE suppress microglial activation and release of TNF.alpha. and NO in a dose-dependent fashion. Peptides derived from the apoE receptor-binding region mimic the effects of the intact protein, whereas deletion of apoE residues 146-149 abolishes peptide bioactivity. These results are consistent with the hypothesis that apoE modulates microglial function by binding specific cell surface receptors and that the immunomodulatory effects of apoE in the central nervous system may account for its role in acute and chronic neurol. disease. (c) 2001 Academic Press.

ST immunomodulator apolipoproteinE Alzheimers disease TNF NO

L8 ANSWER 9 OF 16 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:227511 CAPLUS

DOCUMENT NUMBER:

132:260696

TITLE:

Use of TNF-.alpha. inhibitors for

treating nerve root injury

INVENTOR(S):

Olmarker, Kjell; Rydevik, Bjorn A+ Science Invest AB, Swed.

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 29 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			KIND DATE				APPLICATION NO.					DATE					
											- - ·						
WO	0 2000018409		A1 2		20000406			WO 1999-SE1671					1999				
	W:	ΑE,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,
		CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,
														LT,			
														SE,			
		SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,	AZ,
		BY,	KG,	KZ,	MD,	RU,	ТJ,	TM									
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,
		DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,
		CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG				
SE 9803710			A 20000326					SE 1998-3710				19981029					

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20000417 AU 1999-64918 19990923
    AU 9964918
                      A1
                           20010718 EP 1999-952857 19990923
                      A1
    EP 1115405
        R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
                                      US 2001-760810
                                                           20010117
                      A1 20011004
    US 2001027199
    US 2001027175 A1
                           20011004
                                          US 2001-760811
                                                           20010117
                                       SE 1998-3276 A 19980925
PRIORITY APPLN. INFO.:
                                       SE 1998-3710 A 19981029
                                       WO 1999-SE1671 W 19990923
REFERENCE COUNT:
                        8
                        (2) Olmarker, K; SPINE 1994, V19(16), P1803 MEDLINE
REFERENCE(S):
                         (3) Olmarker, K; SPINE 1998, V23(23), P2538 MEDLINE
                         (4) Pennica, D; NEURON 1996, V17(1), P63 CAPLUS
                         (7) Sommer, C; NEUROSCIENCE LETTERS 1997, V237(1), P45
                            CAPLUS
                         (8) Sommer, C; PAIN 1998, V74(1), P83 CAPLUS
                        ALL CITATIONS AVAILABLE IN THE RE FORMAT
    Use of TNF-.alpha. inhibitors for treating
TI
    nerve root injury
    Pharmaceutical compns. for the treatment of spinal disorders caused by the
AB
    liberation of TNF-.alpha. comprise an effective amt. of a TNF
    -.alpha. inhibitor. Also provided are a method for treatment of
     such disorders and the use of TNF-.alpha. inhibitors
    in the prepn. of a pharmaceutical compn. for such treatment.
    Corticosteroids, biological studies
IT
    Hydroxamic acids
    Lactoferrins
    Tetracyclines
    RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (TNF-.alpha. inhibitors for treating nerve
       root injury)
    Interleukin 1
IT
    RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
        (TNF-.alpha. inhibitors for treating nerve
       root injury)
    Tumor necrosis factors
IT
    RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
     (Biological study); PROC (Process)
        (TNF-.alpha. inhibitors for treating nerve
       root injury)
IT
    Cyclic compounds
    RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (carbocyclic acids; TNF-.alpha. inhibitors for
       treating nerve root injury)
    Carboxylic acids, biological studies
IT
    RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (carbocyclic; TNF-.alpha. inhibitors for treating
       nerve root injury)
    Spinal cord
IT
        (disease; TNF-.alpha. inhibitors for treating
       nerve root injury)
    Nerve, disease
IT
        (injury; TNF-.alpha. inhibitors for treating
       nerve root injury)
    Spinal column
IT
        (intervertebral disk, spinal disk TNF-.alpha.; TNF
       -.alpha. inhibitors for treating nerve root
       injury)
    Steroids, biological studies
IT
    RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (lazaroids; TNF-.alpha. inhibitors for treating
       nerve root injury)
    Spinal column
IT
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(lumbar, nucleus pulposus cells; TNF-.alpha.
       inhibitors for treating nerve root
       injury)
IT
    Antibodies
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (monoclonal, to TNF-.alpha.; TNF-.alpha.
        inhibitors for treating nerve root
        injury)
     Cytokine receptors
IT
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (sol.; TNF-.alpha. inhibitors for treating
       nerve root injury)
     Interferons
IT
     RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
        (.gamma.; TNF-.alpha. inhibitors for treating
       nerve root injury)
                           60-54-8, Tetracycline 60-54-8D, Tetracycline,
     50-35-1, Thalidomide
IT
              73-31-4, Melatonin 79-57-2, Oxytetracycline
                                                               564-25-0,
                   992-21-2, Lymecycline 2444-65-7 10118-90-8, Minocycline
     Doxycycline
     60719-84-8, Amrinone 70458-92-3, Pefloxacin 70458-96-7, Norfloxacin
     74150-27-9, Pimobendan 81840-15-5, Vesnarinone 82419-36-1, Ofloxacin
     85721-33-1, Ciprofloxacin 98079-51-7, Lomefloxacin 108319-06-8,
     Temafloxacin 112811-59-3, Gatifloxacin 170277-31-3, Infliximab
     185243-69-0, Etanercept
     RL: BAC (Biological activity or effector, except adverse); THU
     (Therapeutic use); BIOL (Biological study); USES (Uses)
        (TNF-.alpha. inhibitors for treating nerve
        root injury)
     10102-43-9, Nitrogen oxide (NO), biological studies
IT
     RL: BPR (Biological process); BIOL (Biological study); PROC (Process)
        (TNF-.alpha. inhibitors for treating nerve
        root injury)
     9036-21-9, Phosphodiesterase III 81669-70-7, Metalloproteinase
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (inhibitors; TNF-.alpha. inhibitors for
        treating nerve root injury)
    ANSWER 10 OF 16
                     CAPLUS COPYRIGHT 2001 ACS
                                                       DUPLICATE 3
\mathbf{L8}
ACCESSION NUMBER:
                         2001:9695 CAPLUS
                         134:177129
DOCUMENT NUMBER:
                         Increased production of tumor necrosis factor-.alpha.
TITLE:
                         by glial cells exposed to simulated ischemia or
                         elevated hydrostatic pressure induces apoptosis in
                         cocultured retinal ganglion cells
                         Tezel, Gulgun; Wax, Martin B.
AUTHOR(S):
                         Department of Ophthalmology and Visual Sciences,
CORPORATE SOURCE:
                         Washington University School of Medicine, St. Louis,
                         MO, 63110, USA
                         J. Neurosci. (2000), 20(23), 8693-8700
SOURCE:
                         CODEN: JNRSDS; ISSN: 0270-6474
                         Society for Neuroscience
PUBLISHER:
                         Journal
DOCUMENT TYPE:
                         English
LANGUAGE:
REFERENCE COUNT:
                         66
                         (1) Anderson, D; Invest Ophthalmol Vis Sci 1974, V13,
REFERENCE(S):
                             P771 CAPLUS
                         (2) Barone, F; Stroke 1997, V28, P1233 CAPLUS
                         (4) Bredt, D; Annu Rev Biochem 1994, V63, P175 CAPLUS
                         (5) Brenner, T; Brain Res 1993, V608, P273 CAPLUS
                         (6) Brewer, G; J Neurosci Res 1993, V35, P567 CAPLUS
                         ALL CITATIONS AVAILABLE IN THE RE FORMAT
     Although glial cells in the optic nerve head undergo a reactivation
AB
     process in glaucoma, the role of glial cells during glaucomatous
     neurodegeneration of retinal ganglion cells is unknown. Using a coculture
     system in which retinal ganglion cells and glial cells are grown on
```

'different layers but share the same culture medium, we studied the influences of glial cells on survival of retinal ganglion cells after exposure to different stress conditions typified by simulated ischemia and elevated hydrostatic pressure. After the exposure to these stressors, we obsd. that glial cells secreted tumor necrosis factor-.alpha. (TNF -.alpha.) as well as other noxious agents such as nitric oxide into the coculture media and facilitated the apoptotic death of retinal ganglion cells as assessed by morphol., terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling, and caspase activity. glial origin of these noxious effects was confirmed by passive transfer expts. Furthermore, retinal ganglion cell apoptosis was attenuated .apprx.66% by a neutralizing antibody against TNF-.alpha. and 50% by a selective inhibitor of inducible nitric oxide synthase (1400W). Because elevated intraocular pressure and ischemia are two prominent stress factors identified in the eyes of patients with glaucoma, these findings reveal a novel glia-initiated pathogenic mechanism for retinal ganglion cell death in glaucoma. In addn., these findings suggest that the inhibition of TNF-.alpha. that is released by reactivated glial cells may provide a novel therapeutic target for neuroprotection in the treatment of glaucomatous optic neuropathy.

L8 ANSWER 11 OF 16 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:66204 CAPLUS

DOCUMENT NUMBER:

134:221830

TITLE:

Beneficial effect(s) of n-3 fatty acids in

cardiovascular diseases: but, why and how?

AUTHOR(S):

Das, U. N.

CORPORATE SOURCE:

EFA Sciences LLC, Norwood, MA, 02062, USA

SOURCE:

Prostaglandins, Leukotrienes Essent. Fatty Acids

(2000), 63(6), 351-362

CODEN: PLEAEU; ISSN: 0952-3278

PUBLISHER:

Churchill Livingstone

DOCUMENT TYPE:

Journal; General Review

LANGUAGE:

English

REFERENCE COUNT:

143

REFERENCE(S):

- (4) Besedovsky, H; Science 1986, V233, P652 CAPLUS
- (5) Blann, A; Inflammation 1998, V22, P483 CAPLUS
- (6) Bordet, J; Biochem Biophys Res Commun 1986, V135, P403 CAPLUS
- (7) Bordet, J; Biochim Biophys Acta 1988, V958, P460 CAPLUS
- (8) Borovikova, L; Nature 2000, V405, P458 CAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT
- A review with 143 refs. Low rates of coronary heart disease were found in AB Greenland Eskimos and Japanese who eat diets rich in fish oil. Suggested mechanisms for this cardio-protective effects focused on the effects of n-3 fatty acids on eicosanoid metab., inflammation, fatty acid .beta.-oxidn., endothelial dysfunction, cytokine growth factors, and gene expression of adhesion mols. None of these mechanisms could adequately explain the beneficial actions of n-3 fatty acids. One attractive suggestion is a direct cardiac effect of n-3 fatty acids on arrhythmogenesis. The n-3 fatty acids can modify Na+ channels by directly binding to the channel proteins and thus prevent ischemia-induced ventricular fibrillation and sudden cardiac death. Though this is an attractive explanation, there could be other actions as well. The n-3 fatty acids can inhibit the synthesis and release of proinflammatory cytokines, such as tumor necrosis factor .alpha. (TNF.alpha.) and interleukin-1 (IL-1) and IL-2 released in early ischemic heart disease. These cytokines decrease myocardial contractility, induce myocardial damage, and enhance the prodn. of free radicals which can also suppress myocardial functions. The n-3 fatty acids can increase the parasympathetic tone leading to increased heart rate variability and protection of the myocardium against ventricular arrhythmias. Increased parasympathetic tone and acetylcholine, the principle vagal neurotransmitter, attenuate the release of TNF.alpha., IL-1.beta., IL-6, and IL-18. Exercise enhances the parasympathetic tone

and the prodn. of antiinflammatory cytokine IL-10; this may explain the beneficial action of exercise in the prevention of cardiovascular diseases and diabetes mellitus. TNF.alpha. has neurotoxic actions, whereas n-3 fatty acids are potent neuroprotectors and the brain is rich in these fatty acids. The principal mechanism of the cardioprotective and neuroprotective action(s) of n-3 fatty acids may be due to the suppression of TNF.alpha. and IL synthesis and release, modulation of hypothalamic-pituitary-adrenal antiinflammatory responses, and increased acetylcholine release. There may be close interactions of the central nervous system, endocrine organs, cytokines, exercise, and dietary n-3 fatty acids. This may explain why these fatty acids could be of benefit in the management of conditions such as septicemia and septic shock, Alzheimer disease, Parkinson disease, inflammatory bowel diseases, diabetes mellitus, essential hypertension, and atherosclerosis.

L8 ANSWER 12 OF 16 CAPLUS COPYRIGHT 2001 ACS DUPLICATE 4

ACCESSION NUMBER: 1999:654588 CAPLUS

DOCUMENT NUMBER: 132:120996

TITLE: Intracerebral production of tumor necrosis

factor-.alpha., a local neuroprotective agent, in

Alzheimer disease and vascular dementia

AUTHOR(S): Tarkowski, Elisabeth; Blennow, Kaj; Wallin, Anders;

Tarkowski, Andrzej

CORPORATE SOURCE: Department of Rheumatology, University of Goteborg and

Hospital of Varberg, Swed.

SOURCE: J. Clin. Immunol. (1999), 19(4), 223-230

CODEN: JCIMDO; ISSN: 0271-9142

PUBLISHER: Kluwer Academic/Plenum Publishers

DOCUMENT TYPE: Journal LANGUAGE: English

REFERENCE COUNT: 44

REFERENCE(S): (1) Aarden, L; Eur J Immunol 1987, V17, P1411 CAPLUS

(2) Allsopp, T; Cell 1993, V73, P295 CAPLUS

(4) Anderson, A; J Neurosci 1996, V16, P1710 CAPLUS

(6) Barger, S; Proc Natl Acad Sci USA 1995, V92, P9328 CAPLUS

(11) Brenneman, D; J Neurochem 1992, V58, P454 CAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT

The local pattern of pro-inflammatory cytokine release was studied in Alzheimer disease (AD) and vascular dementia (VAD), by measuring intrathecal levels of IL-1.beta., IL-6, TNF-.alpha., and its naturally occurring antagonists, sol. TNF receptors I and II. The cytokine levels were related to neuronal damage, as measured by the intrathecal tau concn., to cerebral apoptosis assessed by levels of Fas/APO-1 and bcl-2, and to clin. variables. In vitro anal. was performed to study the effect of TNF-.alpha. on the prodn. of bcl-2, an anti-apoptotic factor, by human neuronal cells. Patients with both AD and VAD displayed significantly higher intrathecal levels of TNF-.alpha. compared to controls. In addn., patients with AD showed significantly neg. correlations between the intrathecal levels of TNF-.alpha. and the levels of Fas/APO-1 as well as of tau protein. The level of bcl-2 in supernatants of TNF-.alpha.-exposed cultures of human neuronal cells was up to three times higher than in control supernatants. Our study demonstrates intrathecal prodn. of TNF-.alpha. in patients with dementias, suggesting that this cytokine may have a neuroprotective role in these neurodegenerative conditions as evidenced by neg. correlations between this cytokine and (i) levels of intrathecal Fas/APO-1 and (ii) levels of tau protein, both parameters closely related to brain damage. Our in vitro data suggest that TNF-.alpha. exerts its neuroprotective effect by stimulating neuronal cells to express bcl-2, a mol. which down-regulates apoptosis.

L8 ANSWER 13 OF 16 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V. ACCESSION NUMBER: 1999109361 EMBASE

TITLE: Inhibitory effects of indomethacin on interleukin-1 and

nitric oxide production in rat microglia in vitro.

AUTHOR: Du Z.-Y.; Li X.-Y.

CORPORATE SOURCE: X.Y. Li, Shanghai Institute of Materia Medica, Chinese

Academy of Sciences, Shanghai 200031, China.

xyli@server.shcnc.ac.cn

SOURCE: International Journal of Immunopharmacology, (1999) 21/3

(219-225). Refs: 22

ISSN: 0192-0561 CODEN: IJIMDS

PUBLISHER IDENT.: S 0192-0561(98)00084-8

COUNTRY: United Kingdom
DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 026 Immunology, Serology and Transplantation

030 Pharmacology

037 Drug Literature Index

OO5 General Pathology and Pathological Anatomy

008 Neurology and Neurosurgery

LANGUAGE: English SUMMARY LANGUAGE: English

AB Indomethacin, as a nonsteroidal antiinflammatory drug, is reported to be

effective in some degree in the prevention and treatment of

Alzheimers disease (AD). Effects of indomethacin on proinflammatory cytokines interleukin-1 (IL-1), tumor necrosis factor .alpha. and nitric oxide (NO) on rat microglia. . . IL-1 and NO production by rat microglia stimulated at the concentration of 0.1-10 .mu.mol/1. However, it did not show any inhibitory effect on TNF-.alpha. production by resting and LPS-stimulated rat microglia. The results suggest that the mechanism by which indomethacin might be beneficial in treatment. . .

L8 ANSWER 14 OF 16 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1999:538611 CAPLUS

DOCUMENT NUMBER: 132:77889

TITLE: Downregulation of macrophage activation by PPAR.gamma.

suggests a role for conjugated linoleic acid in

prevention of Alzheimer's disease and atherosclerosis

McCarty, Mark F.

CORPORATE SOURCE: NutriGuard Research, Encinitas, CA, 92024, USA

SOURCE: J. Med. Food (1999), Volume Date 1998, 1(3), 217-226

CODEN: JMFOFJ; ISSN: 1096-620X

PUBLISHER: Mary Ann Liebert, Inc.
DOCUMENT TYPE: Journal; General Review

LANGUAGE: English REFERENCE COUNT: 104

AUTHOR(S):

REFERENCE(S): (1) Allan, C; J Pharmacol Exp Ther 1994, V270, P446 CAPLUS

(2) Altavilla, D; Eur J Pharmacol 1995, V286, P31 CAPLUS

- (3) Angel, P; Biochim Biophys Acta 1991, V1072, P129 CAPLUS
- (4) Bauer, J; Immunol Today 1991, V12, P422 CAPLUS
- (5) Belury, M; Nutr Cancer 1996, V26, P149 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

A review with 104 refs. Activated monocytes/macrophages express the peroxisome proliferator-activated receptor .gamma. (PPAR.gamma.) transcription factor and the activation of PPAR.gamma. with appropriate ligands downregulates the induced macrophage prodn. of interleukin-1 (IL-1) and tumor necrosis factor (TNF). Dietary conjugated linoleic acids (CLA) have thiazolidinedione-like antidiabetic effects in Zucker fatty rats, assocd. with activation of PPAR.gamma. in adipocytes. CLA might exert antiinflammatory effects by suppressing the macrophage cytokine prodn. via PPAR.gamma.. Fish oils rich in n-3 fatty acids also can downregulate the prodn. of IL-1 and TNF by macrophages, possibly because they inhibit autocrine pos. feedback by TXA2. Dietary CLA (fish oil) supplements may be protective with respect to pathologies in which IL-1 and TNF play key etiol. roles. Such pathologies may include atherogenesis and Alzheimer

disease. Antiatherogenic effects of CLA and fish oil have been obsd. in animal models. With regard to Alzheimer disease, the ability of dietary oils to reach the brain implies that CLA/fish oil may have greater clin. utility than drugs that have limited blood-brain barrier penetrance. Available epidemiol. data are consistent with the possibility that frequent fish ingestion may decrease

the risk of Alzheimer disease. ANSWER 15 OF 16 CAPLUS COPYRIGHT 2001 ACS L8ACCESSION NUMBER: 1996:194719 CAPLUS 124:261623 DOCUMENT NUMBER: Preparation of hydroxyalkylammonium-pyrimidines or TITLE: purines and nucleoside derivatives, useful as inhibitors of inflammatory cytokines Benson, Bradley J.; Chen, Xiannong; Cianciolo, George INVENTOR(S): J.; Diaz, Jose-Luis; Ishaq, Khalid S.; Morris-Natschke, Susan L.; Uhing, Ronald J.; Wong, Henry Macronex, Inc., USA PATENT ASSIGNEE(S): PCT Int. Appl., 54 pp. SOURCE: CODEN: PIXXD2 DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE WO 9535304 A1 19951228 WO 1995-US7896 19950621 W: AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, ES, FI, GB, HU, IS, JP, KP, KR, KZ, LK, LU, LV, MG, MN, MW, MX, NO, NZ, PL, PT,

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RO, RU, SD, SE, SG, SK, TM, UA, UG, US, UZ, VN
           RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
                BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG

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OTHER SOURCE(S): MARPAT 124:261623

hydroxyalkylammonioethoxymethylpyrimidine prepn inhibitor inflammatory cytokine; acyclic nucleoside hydroxyalkylammonioethoxymethylpyrimidine prepn; pyrimidine hydroxyalkylammonioethoxymethyl inhibitor inflammatory cytokine; septic shock treatment hydroxyalkylammonioethoxymethylpyrimidine; cachexia treatment hydroxyalkylammonioethoxymethylpyrimidine; rheumatoid arthritis treatment 2134 hydroxyalkylammonioethoxymethylpyrimidine; inflammatory bowel disease treatment 23145 hydroxyalkylammonioethoxymethyl pyrimidine; multiple sclerosis treatment hydroxyalkylammonioethoxymethylpyrimidine; interleukin IL inhibitor hydroxyalkylammonioethoxymethylpyrimidine; tissue factor hydroxyalkylammonioethoxymethylpyrimidine; tissue factor hydroxyalkylammonioethoxymethylpyrimidine; Alzheimer disease hydroxyalkylammonioethoxymethylpyrimidine; Alzheimer

L8 ANSWER 16 OF 16 CAPLUS COPYRIGHT 2001 ACS DUPLICATE 5

ACCESSION NUMBER: 1995:841676 CAPLUS

DOCUMENT NUMBER: 123:254207

TITLE: Tumor necrosis factors .alpha. and .beta. protect

neurons against amyloid .beta.-peptide toxicity: evidence for involvement of a .kappa.B-binding factor

and attenuation of peroxide and Ca2+ accumulation Barger, Steven W.; Hoerster, Dorothee; Furukawa, Katsutoshi; Goodman, Yadong; Krieglstein, Josef;

Mattson, Mark P.

CORPORATE SOURCE: Sanders-Brown Research Center on Aging, Univ.

Kentucky, Lexington, KY, 40536-0230, USA

SOURCE: Proc. Natl. Acad. Sci. U. S. A. (1995), 92(20),

9328-32

CODEN: PNASA6; ISSN: 0027-8424

DOCUMENT TYPE: Journal

AUTHOR(S):

English LANGUAGE: In Alzheimer disease (AD) the amyloid .beta.-peptide AB (A.beta.) accumulates in plaques in the brain. A.beta. can be neurotoxic by a mechanism involving induction of reactive oxygen species (ROS) and elevation of intracellular free calcium levels ([Ca2+]i). In light of evidence for an inflammatory response in the brain in AD and reports of increased levels of tumor necrosis factor (TNF) in AD brain the authors tested the hypothesis that TNFs affect neuronal vulnerability to A.beta.. A.beta.-(25-35) and A.beta.-(1-40) induced neuronal degeneration in a concn. - and time-dependent manner. Pretreatment of cultures for 24 h with TNF-.beta. or TNF -.alpha. resulted in attenuation of A.beta.-induced neuronal degeneration. Accumulation of peroxides induced in neurons by A.beta. was attenuated in TNF-pretreated cultures, and TNFs protected neurons against iron toxicity, suggesting that TNFs induce antioxidant pathways. The [Ca2+]i response to glutamate (quantified by fura-2 imaging) was markedly potentiated in neurons exposed to A.beta., and this action of A.beta. was suppressed in cultures pretreated with TNFs. Electrophoretic mobility-shift assays demonstrated an induction of a .kappa.B-binding activity in hippocampal cells exposed to TNFs. Exposure of cultures to I.kappa.B (MAD3) antisense oligonucleotides, a manipulation designed to induce NF-.kappa.B, mimicked the protection by TNFs. Thus, TNFs protect hippocampal neurons against A.beta. toxicity by suppressing accumulation of ROS and Ca2+ and .kappa.B-dependent transcription is sufficient to mediate these effects. A modulatory role for TNF in the neurodegenerative process in AD is proposed.

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